



CANCER THERAPY EVALUATION PROGRAM, NATIONAL CANCER INSTITUTE

Clinical Trial Update System v3.0 Quick Reference Guide

JULY 1, 2002

FOR USE WITH THE CLINICAL TRIAL UPDATE SYSTEM (CDUS) WEB APPLICATION



BUILDING THE FUTURE™
TOGETHER

PRODUCED BY CAPITAL TECHNOLOGY INFORMATION SERVICES, INC.

The Clinical Trial Update System v3.0 Quick Reference Guide was prepared for:

Cancer Therapy Evaluation Program (CTEP)

Division of Cancer Treatment and Diagnosis (DCTD)

National Cancer Institute (NCI)

National Institutes of Health (NIH)

By:

Capital Technology Information Services, Inc.

1355 Piccard Drive, Suite 450

Rockville, Maryland 20850-4315

Tel: 301-948-3033

Fax: 301-948-2242

Home Page: <http://www.ctisinc.com>

Under the Information Management and Computer Support Contract
NO2-CM-67245.

The brand names and product names used in this manual are trade names, service marks, trademarks, or registered trademarks of their respective owners. All designations appearing in this document that are known to be Service Marks, Trademarks or Registered Trademarks, have been appropriately capitalized. CTIS, Inc. is not associated with any product or vendor mentioned in this manual.

Contents

Introduction	1
Additional Information.....	1
Getting Started	3
Logging On	3
Common CDUS Features.....	4
Formatting.....	4
Icons.....	4
Buttons	5
Navigation.....	5
Error or Warning Messages	5
The Collections Screen	6
Adding a New Collection Record	7
The CDUS Menu	9
Patient Data	11
Patient Data Entry Screens.....	11
Patient Demographics	11
Accessing the Patient Data Entry Screens	12
Patient Administrative Data.....	14
Baseline Abnormalities	16
Prior Therapies.....	17
Treatment Courses	18
Course Agents	19
Adverse Events.....	21
Responses	23
Late Adverse Events	24
Protocol Data	26
Protocol Data Entry Screens	26
Publications.....	26
Authors	27
Correlative Studies.....	29
Phase I End Points MTD and Phase I End Points DLT	29
Phase I End Points MTD	30
Phase I End Points DLT	31
Trial Comments	32
Submissions and Reports	34
Patient Details Report	34
Submitting the Quarterly Clinical Data Update	35

Introduction

The purpose of this guide is to provide the users of the Clinical Trial Update System (CDUS) Web application with concise instructions for accessing and manipulating the version 3.0 (v3.0) enhancements to the CDUS.

The guide walks the user through the process of accessing a data record and adding new or updating existing data to include with the Quarterly Clinical Data Update.

The Quarterly Clinical Data Update is a record that includes all the data collected from each screen in the CDUS Web application. Once complete, the record is sent to CTEP through the CDUS Web application and loaded into the CTEP database (for more information, see the *CDUS Instructions and Guidelines v3.0* available from the CTEP Web site).

Additional Information

The following resources are available to you at the CDUS page of the CTEP Web site:

CDUS Instructions and Guidelines v3.0

Provides details regarding CDUS reporting requirements and detailed descriptions of data elements. This document includes information about the following:

- *CTEP Smart Loader Approval, Disapproval and Correction Process* (pages 52 – 57).
- *Business Rules* (pages 65 – 69). Business rules are used to validate the entry of appropriate or accurate data prior to being saved in the application.

*Clinical Data Update System (CDUS) v3.0, Notice of Modifications,
May 3, 2002*

Provides a comprehensive description of all enhancements included in the CDUS v3.0. The information contained in this document is particularly valuable to those users who have previous experience working with the CDUS Web application.

CTEP Web Site

The CTEP Web site is found at <http://ctep.cancer.gov/> and is accessed to obtain a wide variety of information.

- The CDUS page of the CTEP Web site is located at <http://ctep.cancer.gov/reporting/cdus.html> and provides a link to the CDUS application, to the documents listed above, and to other documents regarding earlier versions of the CDUS.

NCI CTEP Help Desk

Contact the NCI CTEP Help Desk at ncictephhelp@ctep.nci.nih.gov for questions regarding the technical use of the CDUS v3.0 Web application or for training information.

Note: The CDUS Web application is accessed via the Internet using Microsoft Internet Explorer version 5.0 or higher. Use of other browsers or older versions of Microsoft Internet Explorer may cause errors within the application and/or difficulty in its use.

This quick reference guide assumes that you have a working knowledge of Microsoft Windows and Microsoft Internet Explorer browser.

Getting Started

This section of the guide provides instruction and information about the general use of the system and its common elements.

Logging On

Follow the instructions below to log on to the CDUS application.

1. Double-click the **Internet Explorer** (IE) icon on your desktop.
2. Click **Favorites** or select the **Favorites** menu.
3. Select **CDUS Web v3.0** from your Favorites list.

Note: If the CDUS is not available from Favorites, access the CDUS CTEP page and double click on the application link. Once the CDUS main screen displays, add the application to your Favorites list (see your IE manual or IE Help if you are unfamiliar with the Favorites option in IE).

4. Enter your **User Name** and **Password**.
5. Click **OK**.

The **Protocol Selection** screen is displayed (see Figure 1). The **Protocol Number**, **Title**, **Current Trial Status**, and **Current Trial Status Date** are displayed for each protocol listed.

The **Protocol Number** is displayed as a link (see the **Navigation** section on page 5 more information on links).

6. Click on the Protocol Number link for the protocol you wish to access and continue the data entry process.

Note: Only the protocols of the organization for which you have permission will be displayed. This is determined by your User Name and Password. Contact the NCI CTEP Help Desk if there is a discrepancy with the protocols listed from the Protocol Selection screen.

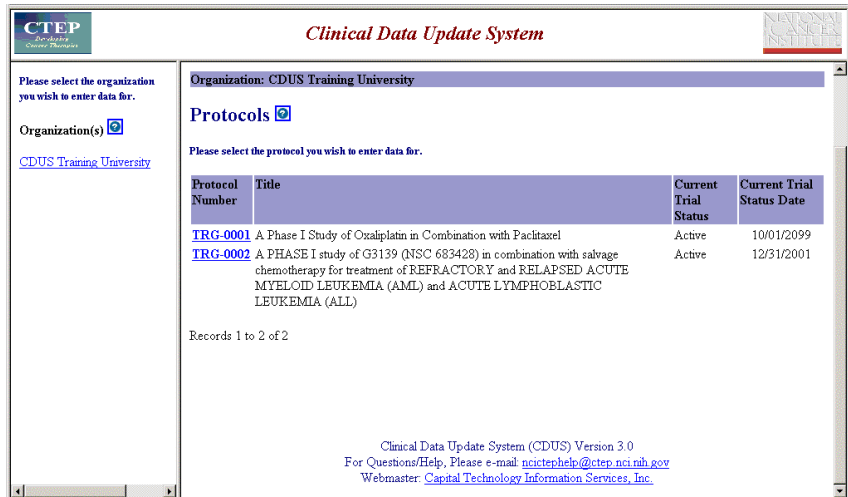


Figure 1: Protocol Selection Screen



Common CDUS Features


Once you have selected a protocol from the Protocol Selections screen, you will find a variety of features that appear throughout the application to assist you in accurately completing the Quarterly Clinical Data Update. The following provides a description of each.


Formatting


Bold Data Elements: Data elements that appear in bold text are mandatory and must be entered prior to clicking the **Save** button. An error message will display when a mandatory data field is left blank.


Icons

 **Protocol Number:** The **Protocol Number** icon is located at the top of each screen and provides access to view a protocol's Organization, Title, Status, and Status Date information. Click on the  to view this information.


 The **Help** icon provides access to view additional instruction and step-by-step processes to assist you while you work with the CDUS. Click the icon to open the Help window.


 The **Calendar** icon is provided as an option for every data element that requires a date and ensures that the date entered is in the correct format. Click the Calendar icon and double click on the day or choose to type the date manually.


 Click the **Up Arrow** icon to the right of a data field to select a value from a List of Values (LOV). Values from the LOV should always be selected, when available, to populate the field.


 Click the **Down Arrow** icon to the right of a data field to select values from a drop-down list.

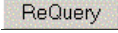
Buttons


 The **Clear** button is available to clear the data from one or all data fields prior to saving.

 The **Delete** button is used to delete a previously saved data record. A message will display prompting you to confirm the delete before the data is removed.

 The **New** button is used to create a new data record. Click the button and a new screen is displayed, from which you will begin data entry.

 The **Query** button is used to search the application for data that matches specified query criteria.

 The **ReQuery** button provides a way to refresh the screen and view a list of data records that were successfully saved.

 The **Save** button is used to commit data to the application. When the data fields are entered correctly and the button is clicked, the message **Success!** is displayed. An error or warning message will display when mandatory data is missing or when an invalid value is entered (see **Error or Warning Messages** on page 5, for additional information).

Navigation

The CDUS Web application uses links to assist you when navigating from one screen to another. Links are presented in blue, underlined text. The links listed on the CDUS menu (see The CDUS Menu section on page 9 for more information) become activated and display the underlined text when the cursor is placed over the screen name.

Error or Warning Messages

The CDUS application uses business rules to validate the entry of appropriate or accurate data. Validations occur each time the **Save** button is clicked, when the **Submit Collections** button is clicked, and again, when the data is loaded to the database at CTEP. Data validations at the screen and submission level may result in an Error and/or Warning message. Data validations that occur during the data load at CTEP may result in an Error Log Report.

The following describes the differences between an Error and Warning message. Again, these messages appear at the time the data is being

saved in any of the CDUS screens or when the Quarterly Clinical Data Update is submitted.

- An Error message is displayed when incomplete or inaccurate data is entered in a **mandatory** data field (see Figure 2). This data must be corrected to commit the data to the application.

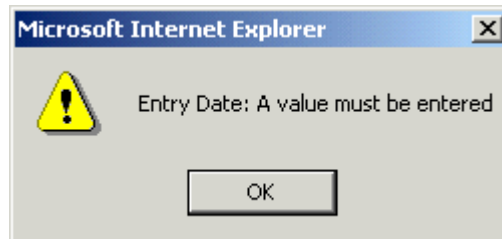


Figure 2: CDUS Error Message

- A Warning message is displayed when incomplete or inaccurate data is entered in a *requested* data field (see Figure 3). Although correction of the data is preferred, it is not mandatory to complete the submission process.

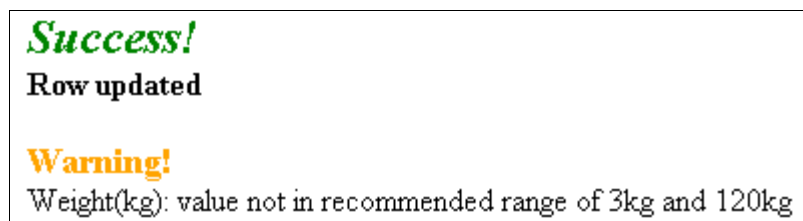


Figure 3: CDUS Warning Message

Follow the instructions below to correct the erroneous data:

1. Click the **OK** button on the CDUS Error Message box or return to the field specified in the Warning Message.
2. Complete, update, or modify the specified data element to correct the error.
3. Click **Save**.

Note: Additional Error or Warning messages may appear if multiple data elements are incomplete or inaccurate. Repeat steps 1 through 3 until all the erroneous data are corrected and no further messages are displayed.

The Collections Screen

To access the **Collections** screen from the **Protocol Selection** screen, click on the Protocol Number link for the protocol you wish to view.

The **Collections** screen is displayed (see Figure 4) and provides a summary of the Quarterly Clinical Data Updates created for present and previous quarters.

Collections

To enter data for a particular collection, please select the collection from the list below. To create a new collection or update an existing collection, select the **Add Collections** button.

	Collection Status	Submission Date (MM/DD/YYYY)	Cutoff Date (MM/DD/YYYY)	Last Submission Date (MM/DD/YYYY)	Current Trial Status	Submitter Name	Submitter Phone	Submitter E-mail
<input type="checkbox"/> Submit?	Active	07/31/2002 (Q2)	06/15/2002		Active	Tamara Whatley	301-402-5924	whatleyt@ma
	Submitted	04/30/2002 (Q1)	03/31/2002	05/22/2002	Active	Betty Boo	3015551555	
	Submitted	01/31/2002 (Q4)	12/31/2001	04/12/2002	Active	Roshini Shank	614-293-4562	shank-2@me
	Accepted	10/31/2001 (Q3)	09/30/2001	12/27/2001	Active	Roshini Shank	6142934562	shank-2@me
	Accepted	07/31/2001 (Q2)	06/30/2001	07/11/2001	Active	Lisa Didier	614-293-8656	didier-1@me
	Accepted	04/30/2001 (Q1)	03/31/2001	04/16/2001	Active	lisa didier	614-293-8656	didier-1@me
	Accepted	01/31/2001 (Q4)	12/31/2000	01/04/2001	Active	lisa didier	614-293-8656	didier-1@me
	Accepted	10/31/2000 (Q3)	09/30/2000	10/10/2000	Active	Lisa Didier	614-293-8656	didier-1@me
	Accepted	07/31/2000 (Q2)	06/30/2000	07/03/2000	Active	Lisa Didier	614-293-8656	didier-1@me
	Accepted	04/30/2000 (Q1)	03/31/2000	04/05/2000	Active	lisa didier	614-293-8656	didier-1@me
	Accepted	01/31/2000 (Q4)	12/31/1999	04/05/2000	Active	Lisa Didier	614-293-8656	didier-1@me

Records 1 to 11 of 11

Note: Active is open for insert and/or update; Submitted and Approved are closed for insert but open for update through the Active collection.

Figure 4: The Collections Screen

The following functions can be performed on the **Collections** screen:

- To enter or update data for an existing Quarterly Clinical Data Update, click on the [Active](#) link from the **Collection Status** column.

Note: Only those Quarterly Clinical Data Update records that appear with an Active or Rejected **Collection Status** may be accessed for new data entry or data update. Records with a status of Submitted, Processing, or Accepted are not available for data entry or update.
- To create a new Quarterly Clinical Data Update or view previously submitted Quarterly Clinical Data Updates, click the **Add Collections** button.
- To submit a completed Quarterly Clinical Data Update, refer to the **Submitting the Quarterly Clinical Data Update** section on page 35.
- To return to the **Protocol Selection** screen, click the **Organization(s)** name listed in the left frame.

Adding a New Collection Record

A new Quarterly Clinical Data Update record must be created for each quarterly data submission. Follow the instructions below to create a new record.

- Click the **Add Collections** button on the **Collections** screen.

The **Collection** data entry screen is displayed (see Figure 5).

Figure 5: The Collection Data Entry Screen

2. Click the **New** button to create a new Quarterly Clinical Data Update record.

Note: The **Submission Date** field is automatically populated with the submission date of the current or subsequent quarter; no data entry is required.

3. Complete all of the mandatory (bold text) data fields and the requested data fields, if relevant information is available.



TIP

The **CutOff Date** field is entered with the latest date for which information is known for this record. The application will validate that all date values entered throughout the remainder of the record will be less than or equal to the Submission Date and less than or equal to the **CutOff Date** identified in this present quarter's record. The present quarter's **CutOff Date** must be greater than or equal to the **CutOff Date** in the previous quarter's record.

4. Click the **Save** button.
5. Click Return to Collection Page link located in the center frame to return to the **Collections** screen.

The new record will display an Active link under the **Collection Status** column. You must click on the Active link to access the CDUS menu where other screens are available to enter and/or update data.

For detailed information regarding the data elements on the **Collection** screen, refer to pages 10 – 12 in the *CDUS Instructions and Guidelines v3.0*.

The CDUS Menu

The CDUS navigation menu resembles a folder directory and lists all of the patient and protocol-specific data entry screens, reports, and navigational links available within the application (see Figure 6).

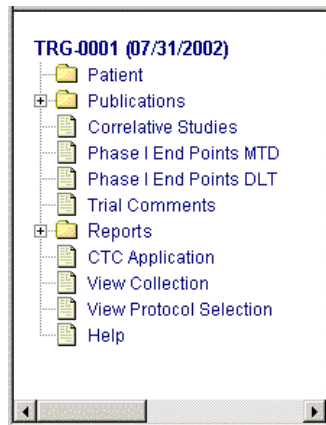


Figure 6: The CDUS Menu

Follow the instructions below to access the CDUS menu.

1. From the **Collections** screen, select a Quarterly Clinical Data Update record by clicking on the **Collection Status Active** link.

The CDUS menu is displayed in the left frame (see Figure 7).

Note: When the **Collection Status Active** link is selected and a patient record exists in the CDUS, the **Patient Demographic Data** screen is displayed by default (as shown in Figure 7).

Patient ID	Birth Date (MM/YYYY)
1	01/1956
2	11/1945
900303191	09/1943
900508297	06/1929
900521553	01/1937
900538668	03/1965
90232	05/1969
906066954	09/1945
906114939	10/1938
906148114	05/1935

Records 1 to 10 of 22

Next Set Last Set

ReQuery

Query

New

Patient Demographic Data

Protocol Number: TRG-0001

Patient ID: 1

Birth Date (MM/YYYY): 01/1956

Gender: Female

Ethnicity: Not Hispanic or Latino

Races: ☒ American Indian or Alaska Native ☐ Asian ☐ Black or African American ☐ Native Hawaiian or Other Pacific Islander ☐ Unknown ☐ White

Country Name: USA

Zip Code: 14569

Payment Method:

Entry Date (MM/DD/YYYY): 01/01/1989

Registering Group:

Registering Institution: A. Maxwell Evans Clinic

Save Delete Clear New

All data elements in bold are mandatory

Figure 7: The CDUS Menu Frame

2. Click on the folder name to view the screen you wish to access.
3. Click on the **+** or **-** sign preceding the folder to expand or collapse a submenu of screens.

If no record exists in the selected screen, only the CDUS menu is displayed in the left frame. The **ReQuery** and **New** buttons are displayed in the center frame. You may click the **New** button to view the data fields available on the selected screen.

If a record was previously entered in the selected screen, the record(s) is listed in the center frame and the first record is displayed in the data entry screen (the right frame) by default.

To return to the **Collections** or the **Protocol Selection** screens, click the View Collection or the View Protocol Selection link from the CDUS menu.

Patient Data

Patient Data Entry Screens

The CDUS Web application provides seven screens to enter patient-specific data and organizes them as follows:

- Demographic Data
- Administrative Data
- Baseline Abnormalities
- Prior Therapies
- Treatment Courses
- Responses
- Late Adverse Events

Note: A new patient demographic record must be created or an existing patient record must be selected from the center frame to access any of the Patient data entry screens. Once a patient is selected, all patient data screens will be specific to the selected patient.

Patient Demographics

The CDUS will provide access to the other patient data entry screens only after the patient demographic record is created. Follow the instructions below to create a new patient demographic record.

Note: Only one Patient Demographic record may be entered per patient.

1. Click on the **Patient** folder from the CDUS menu.

Click the **New** button located in the center frame. A blank demographic data record is displayed in the right frame (see Figure 8).

Figure 8: The Patient Demographic Data Screen

2. Complete all of the mandatory (bold text) data fields and the requested data fields, if relevant information is available.

Select values from the LOV or the drop down list by clicking either the **Up Arrow** or **Down Arrow** buttons.



TIPS

The patient's ethnicity (whether the patient is or is not Hispanic or Latino, or if the patient's ethnicity is unknown) must be indicated within the **Ethnicity** field.

More than one race may be selected from the patient **Races** field.

The **Registering Institution** LOV displays institutions alphabetically by name and includes the CTEP ID, City, State, and Zip code of each. Only the institution name can be used to conduct a search.

Enter the value '00000' if the patient's U.S. Zip code is unknown.

3. Click the **Save** button.

If all data elements are entered correctly, the message **Success! Row inserted** will display in the top left of the screen. If a mandatory data field was missed or data were inaccurately entered, an error or warning message will display (see **Error or Warning Messages** on page 5, for additional information).

For detailed information regarding the Patient Demographic data elements, refer to pages 19 – 23 in the [CDUS Instructions and Guidelines v3.0](#).

Accessing the Patient Data Entry Screens

Once the patient demographic record is saved, the **Patient ID** and **Birth Date** are displayed in the center frame (see Figure 9). The

Patient ID entered in the **Patient Demographic Data** screen is displayed as a link under the **Patient ID** column. You must click on the Patient ID link to make modifications in **Patient Demographic Data** screen or to access other patient data entry screens.

The screenshot shows a window titled "Patients". Inside, there is a prompt "Please select a patient to proceed". Below this is a table with two columns: "Patient ID" and "Birth Date (MM/YYYY)". The first row contains the values "90232" and "05/1969". Below the table, it says "Records 1 of 1". There are several buttons: "Next Set", "Last Set", "ReQuery", "Query", and "New".

Patient ID	Birth Date (MM/YYYY)
90232	05/1969

Figure 9: The Patients Record (center frame)

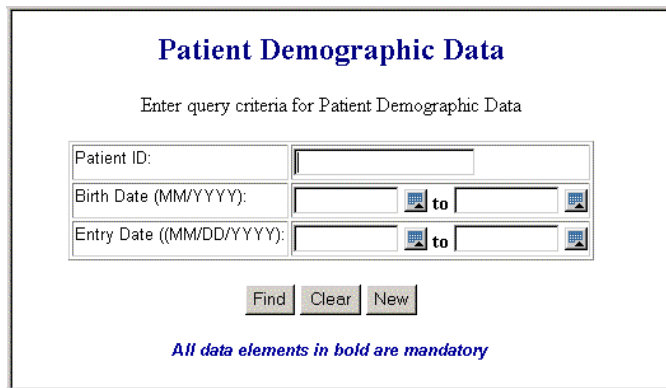
When the Patient ID link is selected for a patient, the Patient folder under the CDUS menu expands to display the screens available for patient data entry (see Figure 10). The Patient ID and Birth Date are also displayed within parentheses following the Patient folder.

The screenshot shows a tree view under the "TRG-0001 (07/31/2002)" menu. The "Patient (90232, 05/1969)" folder is expanded, showing a list of sub-items: Administrative Data, Baseline Abnormalities, Prior Therapies, Treatment Courses, Responses, Late Adverse Events, Publications, Correlative Studies, Phase I End Points MTD, Phase I End Points DLT, Trial Comments, Reports, Patient Details, CTC Application, View Collection, View Protocol Selection, and Help.

Figure 10: The Patient Folder – Expanded

The center frame is not capable of displaying all the Patient ID links associated with a protocol where a large number of patients are enrolled. In this case, a search must be conducted to access the record

of a specific patient. A search can be performed by clicking the **Next Set** and **Last Set** buttons from the center frame or by using the **Patient Demographic Data Query** screen (see Figure 11).



The screenshot shows a web application window titled "Patient Demographic Data". Below the title is the instruction "Enter query criteria for Patient Demographic Data". There are three input fields: "Patient ID:" with a single text box, "Birth Date (MM/YYYY):" with two text boxes separated by a "to" label and calendar icons, and "Entry Date ((MM/DD/YYYY):" with two text boxes separated by a "to" label and calendar icons. Below these fields are three buttons: "Find", "Clear", and "New". At the bottom, a note states "All data elements in bold are mandatory".

Figure 11: Patient Demographic Data Query Screen

To use the **Patient Demographic Data Query** screen, click the **Query** button from the center frame, enter criteria specific to the patient in any of the available fields, and click the **Find** button.

Note: The percentage symbol (%) can be used as a wildcard within the **Patient ID** field only.

Patient Administrative Data

Patient administrative data is mandatory for trials assigned to complete CDUS reporting. Follow the instructions below to enter patient-specific administrative data.

Note: Only one Patient Administrative record may be entered per patient.

1. Click on the Patient ID link located in the center frame under the **Patient ID** column for the patient record you wish to access.
2. Select the Administrative Data link from the CDUS menu. The **Patient Administrative Data** screen is displayed for the selected **Patient ID** in the left frame (see Figure 12).

Figure 12: The Patient Administrative Data Screen

3. Complete all of the mandatory (bold text) data fields and the requested data fields, if relevant information is available.

Select values from the LOV or the drop down list by clicking either the **Up Arrow** or **Down Arrow** buttons.



TIPS

The **Baseline Abnormalities** screen must be completed if 'Yes' is entered in the **Has the Patient had any Baseline Abnormalities?** field (see the Baseline Abnormalities section on page 16 for more information).

The **Off Treatment Reason** field becomes mandatory if 'No' is entered in the **Is the Patient currently receiving treatment on study?** field.

The **Last Treatment Date** field becomes mandatory when the **Off Treatment Reason** field is entered. This rule does not apply when an **Off Treatment Reason** value of 'Patient withdrawal before beginning Active Treatment' or 'Disease Progression before Active Treatment' is entered.

Note: The term *Active Treatment* is considered any form of therapy (including surgery, radiation, commercial chemotherapy agents or investigational agents).

The **Off Study Reason** field becomes mandatory when the **Off Study Date** field is entered.

4. Click the **Save** button.

For detailed information regarding the Patient Administrative data elements, refer to pages 23 - 28 in the [CDUS Instructions and Guidelines v3.0](#).

Baseline Abnormalities

The **Baseline Abnormalities** screen is mandatory if you indicated that the patient had baseline abnormalities in the **Patient Administrative Data** screen. Follow the instructions below to enter baseline abnormalities for a selected patient.

Note: Multiple Baseline Abnormality records may be entered per patient.

1. Select Baseline Abnormalities from the CDUS menu.
2. Click the **New** button. The **Baseline Abnormalities** screen is displayed in the right frame (see Figure 13).

TRG-0001 (07/31/2002)

Patient (90232, 05/1969)

- Administrative Data
- Baseline Abnormalities
- Prior Therapies
- Treatment Courses
- Responses
- Late Adverse Events
- Publications
- Correlative Studies
- Phase I End Points MTD
- Phase I End Points DLT
- Trial Comments
- Reports
- CTC Application
- View Collection
- View Protocol Selection
- Help

Baseline Abnormalities

No Records returned

ReQuery

New

Protocol Number: TRG-0001

Baseline Abnormalities

Patient ID: 90232

Birth Date: 05/1969

Enter values for new Baseline Abnormalities record

Category:

Adverse Event:

Other Adverse Event (Specify):

Grade:

Save Clear

All data elements in bold are mandatory

Figure 13: The Baseline Abnormalities Screen

3. Complete all of the mandatory (bold text) data fields and the requested data fields, if relevant information is available.

Select values from the LOV by clicking the **Up Arrow** button.



TIP

If you select 'Other Specify' for the **Adverse Event** field, you must provide the specific Adverse Event in the **Other Adverse Event** field.

4. Click the **Save** button.

The **Category**, **Adverse Event**, and **Grade** of the Baseline Abnormalities record are displayed in the center frame (see Figure 14). If needed, you may click the Category link to access and update the record.

Category	Adverse Event	Grade
CARDIOVASCULAR (GENERAL)	Hypertension	1

Record 1 of 1

Figure 14: The Baseline Abnormalities Record (center frame)

- To enter multiple baseline abnormality records, Click the **New** button and repeat steps 2 through 4 for each record.

For detailed information regarding the Baseline Abnormalities data elements, refer to page 28 in the [CDUS Instructions and Guidelines v3.0](#).

Prior Therapies

Prior therapies are mandatory for trials assigned to complete CDUS reporting. Follow the instructions below to enter all prior cancer therapies the patient has received.

Note: Multiple Prior Therapies records may be entered per patient. Up to five therapies can be entered at one time.

- Select the [Prior Therapies](#) link from the CDUS menu.
- Click the **New** button. The **Prior Therapies** screen is displayed (see Figure 15).

TRG-0001 (07/31/2002)

Protocol Number: TRG-0001

Prior Therapies

Patient ID: 90232
Birth Date: 05/1969

Therapy	Insert?
<input type="text"/>	<input type="button" value="Clear"/>
<input type="text"/>	<input type="button" value="Clear"/>
<input type="text"/>	<input type="button" value="Clear"/>
<input type="text"/>	<input type="button" value="Clear"/>
<input type="text"/>	<input type="button" value="Clear"/>

All data elements in bold are mandatory

Figure 15: The Prior Therapies Screen

- Click the **Down Arrow** button and select a **Prior Therapy** value from the drop down list.

4. Click the **Clear** button if you wish to remove a Prior **Therapy** value from the list.
5. Click the **Save** button. The entered therapies are displayed.
6. To remove any Prior **Therapy** value from the saved list, click the **Delete** checkbox and click the **Save** button.
7. To enter additional prior therapies, click the **New** button and repeat steps 2 through 5 above.

For detailed information regarding Prior Therapies data elements, refer to pages 26 - 28 in the *CDUS Instructions and Guidelines v3.0*.

Treatment Courses

Treatment course data is mandatory for trials assigned to complete CDUS reporting. Follow the instructions below to enter treatment course data.

Note: Multiple Treatment Course records may be entered per patient.

1. Select the Treatment Courses link from the CDUS menu.
2. Click the **New** button. The **Treatment Courses** screen is displayed (see Figure 16).

Figure 16: The Treatment Courses Screen

3. Complete all of the mandatory (bold text) data fields and the requested data fields, if relevant information is available.

Select values from the LOV or the drop down list by clicking either the **Up Arrow** or **Down Arrow** buttons.



TIP

If you enter 'Yes' in the **Adverse Event Experienced** field, you must provide specific Adverse Event data in the **Adverse Event** screen (see **Adverse Events** on page 21).

4. Click the **Save** button.

The **Course ID**, **Course Start Date**, and **Treatment Assignment** of the Treatment Course record are displayed in the center frame (see Figure 17). If needed, you may click the Course ID link to access and update the record.

Course ID	Course Start Date (MM/DD/YYYY)	Treatment Assignment
2	06/23/2002	TAA
1	05/09/2002	TAB

Records 1 to 2 of 2

ReQuery

New

Figure 17: The Treatment Courses Record (center frame)

To complete the Treatment Courses process, you must enter information in the **Course Agents** screen and in the **Adverse Events** screen if 'Yes' was entered in the **Adverse Event Experienced?** field of the **Treatment Courses** screen. Follow the instructions below to complete the data entry process for these screens.

Course Agents

Note: Multiple Course Agent records may be entered per patient.

1. Click the Course ID link in the center frame. The Course Agents and Adverse Events links are displayed on the CDUS menu (see Figure 18).

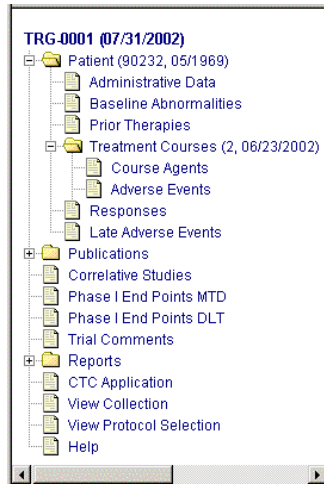


Figure 18: The Course Agents and Adverse Events Links

2. Click the Course Agents link from the CDUS menu.
3. Click the **New** button located in the center frame. The **Course Agents** screen is displayed (see Figure 19). The **Course ID** and **Treatment Assignment** fields in the right frame are automatically populated.

Figure 19: The Course Agents Screen

4. Enter the **Agent Name** field and enter the dose information for the agent the patient received on the selected Treatment Course.
5. Click the **Save** button.

The **Course Agent** is displayed as a link in the center frame (see Figure 20). If needed, you may click the Course Agent link to access and update the record.

Course Agent
OXALIPLATIN
TAXOL

Records 1 to 2 of 2

ReQuery

New

Figure 20: The Course Agents Record (center frame)

6. To enter additional agent records, click the **New** button and follow steps 3 through 5 above.
7. Click the Treatment Courses link on the CDUS menu to return to the **Treatment Courses** screen or click on the Adverse Events link on the CDUS menu to complete the Treatment Courses data entry.

Adverse Events

The **Adverse Event** screen is displayed only when 'Yes' is entered in the **Adverse Event Experienced?** field of the **Treatment Courses** screen.

Note: Multiple Adverse Event records may be entered per patient.

1. Click the Course ID link from the center frame. The Course Agents and Adverse Events links are displayed on the CDUS menu as (see Figure 18).
2. Click the Adverse Events link from the CDUS menu. The **Adverse Event** screen is displayed
3. Click the **New** button located in the center frame. The **Adverse Events** screen is displayed (see Figure 21). The **Course ID** and **Treatment Assignment** fields are automatically populated.

Figure 21: The Adverse Event Screen

4. Complete all of the mandatory (bold text) data fields and the requested data fields, if relevant information is available.

Select values from the LOV or the drop down list by clicking either the **Up Arrow** or **Down Arrow** buttons.



TIP

If you enter 'Other Specify' in the **Adverse Event** field, you must provide the specific Adverse Event in the **Other Adverse Event** field.

5. Click the **Save** button.
6. The **Category**, **Adverse Event**, and **Grade** of the Adverse Event record are displayed as a link in the center frame (see Figure 22). If needed, you may click the Category link to access and update the record.

Category	Adverse Event	Grade
CARDIOVASCULAR (GENERAL)	Hypertension	2

Record 1 of 1

ReQuery

New

Figure 22: The Adverse Event Record (center frame)

7. To enter additional Adverse Event records, click the **New** button and follow steps 3 through 5 above, and enter data for each event.

- Click the Treatment Courses link on the CDUS menu to return to the **Treatment Courses** screen.

For detailed information regarding the Treatment Courses, Course Agents, and Adverse Events data elements, refer to pages 28 - 34 in the CDUS Instructions and Guidelines v3.0.

Responses

Response data is mandatory when 'Yes' is entered in the **Is the Patient Evaluable for Response?** field from the **Administrative Data** screen. Follow the instructions below to enter response data for the selected patient.

Notes: A Treatment Course record must be created prior to entering response information.

Multiple Response records may be entered per patient. Up to five responses can be entered at one time.

- Select the Responses link from the CDUS menu.
- Click the **New** button. The **Responses** screen is displayed (see Figure 23).

Category	Observed Date	Insert?
		Clear
		Clear
		Clear
		Clear
		Clear

Save

All data elements in bold are mandatory

Figure 23: The Responses Screen

- Click the **Down Arrow** button and select a Response **Category** value from the drop down list. Enter the **Observed Date**.



TIPS

Only enter the patient's earliest observed best response.

Progression should be reported even if it is experienced after a better response.

The values entered in the Response **Category** field should not decline except to the value 'Progression.'

Other Response **Category** values will not be accepted if 'Progression' is entered as the initial value.

When 'Other' is entered as the Response **Category** value, the General Response Comments field will be mandatory in the **Trial Comments** (see page 32) screen.

4. Click the **Clear** button if you wish to remove a Response **Category** value from the list.
5. Click the **Save** button. The entered responses are displayed.
6. To remove any Response value from the saved list, click the **Delete** checkbox and click the **Save** button.
7. To enter additional response records, click the **New** button and follow steps 2 through 5 above.

For detailed information regarding Response data elements, refer to pages 35 - 36 in the *CDUS Instructions and Guidelines v3.0*.

Late Adverse Events

Complete the **Late Adverse Event** screen when an Adverse Event is observed after a patient has completed treatment. Follow the instructions below to enter Late Adverse Events.

Note: Multiple Late Adverse Event records may be entered per patient.

1. Select the Late Adverse Events link from the CDUS menu.
2. Click the **New** button from the center frame. The **Late Adverse Event** screen is displayed (see Figure 24).

Figure 24: The Late Adverse Event Screen

3. Complete all of the mandatory (bold text) data fields and the requested data fields, if relevant information is available.
Select values from the LOV by clicking the **Up Arrow** button.



TIP

If you enter 'Other Specify' in the **Adverse Event** field, you must provide the specific Adverse Event in the **Other Adverse Event** field.

4. Click the **Save** button.
5. The **Category**, **Adverse Event**, and **Grade** of the Late Adverse Event record are displayed as a link in the center frame (see Figure 25). If needed, you may click the Category link to access and update the record.

Category	Adverse Events	Grade
CARDIOVASCULAR (GENERAL)	Hypertension	2

Record 1 of 1

Figure 25: The Late Adverse Event Record (center frame)

6. To enter additional Late Adverse Event records, click the **New** button and follow steps 2 through 4 above.

For detailed information regarding Late Adverse Event data elements, refer to pages 34 and 35 in the [CDUS Instructions and Guidelines v3.0](#).

Protocol Data

Protocol Data Entry Screens

The CDUS Web application provides five screens to enter protocol-specific data and organizes them as follows:

- Publications
- Correlative Studies
- Phase I End Points MTD
- Phase I End Points DLT
- Trial Comments

Publications

A publication citation must be provided when data for the study or any associated correlative study is published. Follow the instructions below to enter Publications data.

Note: Multiple Publications records may be entered per protocol.

1. Select the Publications link from the CDUS menu.
2. Click the **New** button from the center frame. The **Publications** screen is displayed (see Figure 26).

TRG-0001 (07/31/2002)

- Patient (90232, 05/1969)
 - Administrative Data
 - Baseline Abnormalities
 - Prior Therapies
 - Treatment Courses
 - Responses
 - Late Adverse Events
- Publications**
- Correlative Studies
- Phase I End Points MTD
- Phase I End Points DLT
- Trial Comments
- Reports
- CTC Application
- View Collection
- View Protocol Selection
- Help

Publications

No Records returned

ReQuery

New

Protocol Number: TRG-0001

Publications

Enter values for new Publications record

Medline UID:

Title:

Journal:

Volume:

Year (YYYY):

Publisher:

Pages:

Save Clear

All data elements in bold are mandatory

Figure 26: The Publications Screen

3. If the publication has an assigned Medline Unique Identifier (UID), you need only enter the **Medline UID** field. If no Medline UID is available, then all other data fields must be entered to complete the Publications record.
4. Click the **Save** button.
5. The **Medline UID** or article **Title** of the Publications record is displayed as a link in the center frame (see Figure 27). If needed, you may click the Medline UID or Title link to access and update the record.

Medline UID	Title
AG223334	Oxaliplatin in Combination with Paclitaxel

Records 1 to 2 of 2

[ReQuery](#)

[New](#)

Figure 27: The Publications Record (center frame)

6. To enter additional Publications records, click the **New** button and follow steps 2 through 4 above.


To complete the Publications process, you must enter information in the **Authors** screen. Follow the instructions below to complete the data entry process for this screen.

Authors

All authors associated with the article should be entered for each Publication record.

Notes: Multiple Author records may be entered per Publication. Up to five author names may be entered at one time.

Author information is not necessary if the Medline UID was entered.

1. On the CDUS menu, click on the  preceding the Publications folder to expand and view the subfolder.
2. Select the Authors link from the CDUS menu.

- Click the Medline UID or the article Title link in the center frame to select the Publication record you wish to add authors to.
- Click the **New** button from the center frame. The **Authors** data entry screen is displayed (see Figure 28).

Figure 28: The Authors Screen

- Enter the Author's last, first, and middle name(s) in the same order as they appear in the selected publication.
- Click the **Clear** button if you wish to remove an Author's name from the list.
- Click the **Save** button. The **Rows inserted successfully** message is displayed.
- To view the entered Authors, click on the Medline UID or the article Title link in the center frame. The Author records are displayed (see Figure 29).

Figure 29: The Authors Record

- To remove any Author name from the saved list, click the **Delete** checkbox and click the **Save** button.
- To enter additional Author names, click the **New** button and repeat steps 4 through 8 above.

For detailed information regarding Publications data elements, refer to pages 18 - 19 in the CDUS Instructions and Guidelines v3.0.

Correlative Studies

Correlative study data must be provided for each correlative study every quarter when correlative studies are associated with the protocol. Follow the instructions below to enter correlative study data.

Notes: A separate Correlative Study record is automatically created for each correlative study associated with the protocol.

Only one Correlative Study record is available per correlative study.

1. Select the Correlative Studies link from the CDUS menu. The **Correlative Studies** screen is displayed (see Figure 30).

Figure 30: The Correlative Studies Screen

2. Click on the Study Code link in the center frame for the Correlative Study record you wish to access.
3. Complete all of the mandatory (bold text) data fields and the requested data field, if relevant information is available.
4. Click the **Save** button.

For detailed information regarding Correlative Studies data elements, refer to page 16 in the CDUS Instructions and Guidelines v3.0.

Phase I End Points MTD and Phase I End Points DLT

Phase I end points include the recommended Phase 2 dose or maximum tolerated dose (MTD) and dose limiting toxicity (DLT) information. This information is mandatory for Phase I studies assigned to complete CDUS reporting.

The Phase I End Points MTD and DLT are identified by the subgroup and treatment assignment for which the DLT occurred and the MTD determined. This data combination creates a unique data key, which assists CTEP in further understanding the agent's abilities. The MTD and DLT information is expected towards the completion of the trial.

Follow the instructions below to enter Phase I End Points data.

Phase I End Points MTD

Note: Multiple Phase I End Points MTD records may be entered per protocol.

1. Select the Phase I End Points MTD link from the CDUS menu.
2. Click the **New** button. The **Phase I End Points MTD** screen is displayed (see Figure 31).

Figure 31: The **Phase I End Points MTD** Screen

3. Complete all of the mandatory (bold text) data fields by clicking the **Up Arrow** button and entering values from the LOV.
4. Click the **Save** button.
5. The **Subgroup Code** and **Treatment Assignment** of the **Phase I End Points MTD** record are displayed as a link in the center frame (see Figure 32). If needed, you may click the Subgroup Code link to access and update the record.

Figure 32: The **Phase I End Points MTD** Record (center frame)

6. To enter additional Phase I End Points MTD records, click the **New** button and follow steps 2 through 4 above.

Phase I End Points DLT

Note: Multiple Phase I End Points DLT records may be entered per protocol.

1. Select the Phase I End Points DLT link from the CDUS menu.
2. Click the **New** button. The **Phase I End Points DLT** screen is displayed (see Figure 33).

Figure 33: The Phase I End Points DLT Screen

3. Complete all of the mandatory (bold text) data fields by clicking the **Up Arrow** button and entering values from the LOV.
4. Click the **Save** button.
5. The **Subgroup Code**, **Treatment Assignment**, and **Adverse Event** of the Phase I End Points DLT record are displayed as a link in the center frame (see Figure 34). If needed, you may click the Subgroup Code link to access and update the record.

Figure 34: The Phase I End Points DLT Record (center frame)

6. To enter additional Phase I End Points DLT records, click the **New** button and follow steps 2 through 4 above.

Trial Comments

The **Trial Comments** screen is used to provide a general data summary by subgroup and treatment assignment. This screen is optional. Follow the instructions below to enter Trial Comments data.

1. Click the [Trial Comments](#) link from the CDUS menu.
2. Click the **New** button. The **Trial Comments** screen is displayed (see Figure 35).

TRG-0001 (07/31/2002)

- Patient
- Publications
- Correlative Studies
- Phase I End Points MTD
- Phase I End Points DLT
- Trial Comments
- Reports
- CTC Application
- View Collection
- View Protocol Selection
- Help

Trial Comments

No Records returned

ReQuery

New

Protocol Number: TRG-0001

Trial Comments

Enter values for new Trial Comments record

Subgroup Code:

Treatment Assignment:

General Response Comments:

General Adverse Events Comments:

Save Clear

All data elements in bold are mandatory

Figure 35: The Trial Comments Screen

3. Complete all of the mandatory (bold text) data fields by clicking the **Up Arrow** button and entering values from the LOV. Complete the requested data fields, if relevant information is available.



TIP

When 'Other' is entered as the Response **Category** value in the **Responses** screen (see page 23), the General Response Comments field will be mandatory in the **Trial Comments** screen.

4. Click the **Save** button.
5. The **Subgroup Code** and **Treatment Assignment** of the Trial Comments record are displayed as a link in the center frame (see Figure 36). If needed, you may click the [Subgroup Code](#) link to access and update the record.

The screenshot shows a web application window titled "Trial Comments". Inside, there is a table with two columns: "Subgroup Code" and "Treatment Assignment". The "Subgroup Code" column contains the text "SG1" and the "Treatment Assignment" column contains the text "TAB". Below the table, it says "Record 1 of 1". At the bottom of the form, there are two buttons: "ReQuery" and "New".

Subgroup Code	Treatment Assignment
SG1	TAB

Record 1 of 1

ReQuery

New

Figure 36: The Trial Comments Record (center frame)


6. To enter additional Trial Comments records, click the **New** button and follow steps 2 through 4 above.

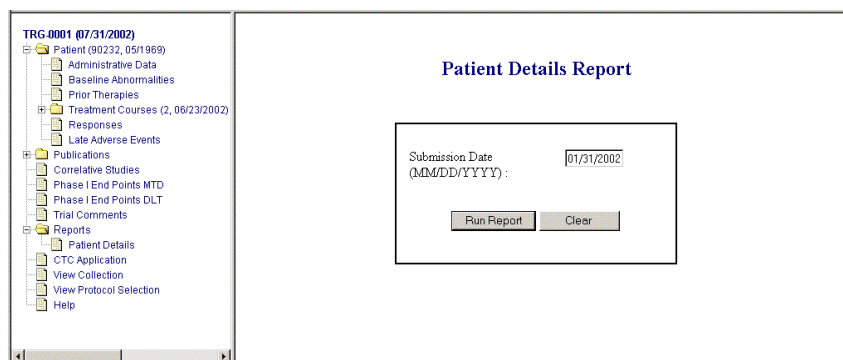
For detailed information regarding Trial Comments data elements, refer to page 18 in the *CDUS Instructions and Guidelines v3.0*.

Submissions and Reports

Patient Details Report

The **Patient Details Report** provides the current data for each patient enrolled on the protocol and entered in the Quarterly Clinical Data Update. Because the report is cumulative, it includes all new patient records entered for the quarter and any modifications made to existing patient data. The report does not show original or previous data once the data is modified. Follow the instructions below to generate the **Patient Details Report**.

1. On the CDUS menu, click on the  preceding the Reports folder to expand and view the subfolders.
2. Select the Patient Details link from the CDUS menu. The **Patient Details Report** generation screen is displayed (see Figure 37).



TRG-0001 (07/31/2002)

- Patient (90232, 05/1969)
 - Administrative Data
 - Baseline Abnormalities
 - Prior Therapies
 - Treatment Courses (2, 06/23/2002)
 - Responses
 - Late Adverse Events
- Publications
- Correlative Studies
- Phase I End Points MTD
- Phase I End Points DLT
- Trial Comments
- Reports
 - Patient Details**
 - CTC Application
 - View Collection
 - View Protocol Selection
 - Help

Patient Details Report

Submission Date (MM/DD/YYYY) : 01/31/2002

Figure 37: The Patient Details Report Generation Screen

3. Enter the date that the Quarterly Clinical Data Update was submitted in the **Submission Date** field.
4. Click the **Run Report** button.

The **Patient Details Report** is displayed as an Adobe Acrobat PDF file (see Figure 38).

Clinical Data Update System Patient Details Report as of 01/31/2001						Page 1 of 12	
Protocol Number: TRG-0001		Run By: CDUS User (06/17/2002)					
Title: A Phase I Study of Oxaliplatin in Combination with Paclitaxel							
Patient ID:	90252	Birth Date:	05/1969	Entry Date:	05/22/2002		
Gender:	Female	Ethnicity:	Not Hispanic or Latino	Country:	USA		
Registering Group:				Registering Institution:	NCIMET-National Cancer Institute Metabolism Branch		
Subgroup:	SG1 - Patients with refractory solid tumors						
Disease:	Leukoplakia esophageal						
Has the patient been declared ineligible?:	No	Is the patient evaluable for Response?:	Yes				
Is the patient currently receiving treatment on study?:	Yes	Off Treatment Reason:					
Last Treatment Date:		Off Study Reason & Date:	-				
Number of Prior Chemotherapy Regimens:	2	Baseline Abnormality Flag:	Yes				
Race:	Native Hawaiian or Other Pacific Islander						
Patient Response:	No Patient Responses Reported						
Prior Therapy:	No Prior Therapy Reported						
Baseline Abnormalities Category:	Adverse Event:	Grade:	AE Other Specific:				
CARDIOVASCULAR (General)	Hypertension	1					
Treatment Course:	Course Start Date:	Treatment Assignment:	Treating Institution:	Height (cm):	Weight (kg):	AE Experienced?	
1	05/09/2002	TAB	OH007 CDUS Training	190.5	83.5	No	
2	06/23/2002	TAA					
Late Adverse Event:	No Late Adverse Events Reported						

Figure 38: The Patient Details Report

The report uses the assigned Patient ID to organize the report data and displays the patient records in alphanumeric order.

Submitting the Quarterly Clinical Data Update

Once all the data are entered for the Quarterly Clinical Data Update, it is submitted to the Cancer Therapy Evaluation Program (CTEP). Follow the instructions below to submit the Quarterly Clinical Data Update.

1. Update and/or enter new data in all Patient and Protocol screens. Review the data for accuracy.
2. Select the protocol to access the **Collections** screen.
3. Check the **Submit?** checkbox located in the first column of the table.
4. Click the **Submit Collections** button.

The Quarterly Clinical Data Update is now submitted to CTEP. If an Error Message is displayed, correct the error by following the instructions provided in the **Error or Warning Messages** section on page 5.